## Amendments to the Claims:

This listing of claims will replace all prior versions, and listings of claims in the application:

## **Listing of Claims**

- 1. (Currently Amended) A method for the treatment of a <u>renal cell</u> cancer comprising co-administering an anti-tumor antibody directed against the MN antigen and a cytokine to a subject in need thereof, wherein the cytokine is <u>an interferon and is</u> administered continuously or repeatedly in a low-dose form, <u>wherein the low-dose cytokine comprises a dose which is pharmaceutically</u> effective in the absence of NIC CTC toxicity grade 3 or higher.
- 2. (Currently Amended) A method for the treatment of a <u>renal cell</u> cancer comprising co-administering an anti-tumor antibody directed against the MN antigen and cytokine to a subject in need thereof, wherein the <u>cytokine is an interferon and the</u> method comprises:
- (a) a first treatment stage comprising administering a low-dose cytokine, and
- (b) a second treatment stage comprising co-administering the anti-tumor antibody and a low-dose cytokine, and wherein the low-dose cytokine comprises a dose which is pharmaceutically effective in the absence of NIC CTC toxicity grade 3 or higher.
- 3. (Cancelled)

4.	(Previously Presented) The method according to claim 1 comprising a			
daily administration of a low-dose cytokine.				
5.	(Cancelled)			
6.	(Cancelled)			
7.	(Cancelled)			
8.	(Currently Amended) The method of claim $\frac{5}{1}$ wherein the cytokine is			
IFN-α				
9.	(Original) The method of claim 8 wherein the dose of IFN- $\alpha$ is in the range			
of from 1-10 MIU three times a week.				
10.	(Currently Amended) The method of claim 1 wherein the cytokine is			
administered in a substantially constant dose during the treatment.				
11.	(Previously Presented) The method of claim 1 wherein the cytokine is			
administered in a variable dose during the treatment.				

12.	(Previously Presented)	The method of claim 1	wherein the cytokine is
admir	nistered subcutaneously.		

## 13. (Cancelled)

- 14. (Previously Presented) The method of claim 1 wherein the antitumor antibody is a chimeric or humanized G250 antibody or a fragment thereof.
- 15. (Previously Presented) The method of claim 1 wherein the antitumor antibody is administered in intervals of from 5-20 days.
- 16. (Original) The method of claim 2 wherein the first treatment stage comprises 5-20 days.
- 17. (Original) The method of claim 2 wherein the second treatment stage comprises 50-200 days.